§610.67

§ 610.67 Bar code label requirements.

Biological products must comply with the bar code requirements at §201.25 of this chapter. However, the bar code requirements do not apply to devices regulated by the Center for Biologics Evaluation and Research or to blood and blood components intended for transfusion. For blood and blood components intended for transfusion, the requirements at §606.121(c)(13) of this chapter apply instead.

[69 FR 9171, Feb. 26, 2004]

PART 630—GENERAL REQUIRE-MENTS FOR BLOOD, BLOOD COMPONENTS, AND BLOOD DE-RIVATIVES

AUTHORITY: 21 U.S.C. 321, 331, 351, 352, 355, 360, 371; 42 U.S.C. 216, 262, 264.

SOURCE: 66 FR 31176, June 11, 2001, unless otherwise noted.

§630.6 Donor notification.

- (a) Notification of donors. You, an establishment that collects blood or blood components, must make reasonable attempts to notify any donor, including an autologous donor, who has been deferred based on the results of tests for evidence of infection with a communicable disease agent(s) as required by §610.41 of this chapter; or who has been determined not to be suitable as a donor based on suitability criteria under §640.3 or §640.63 of this chapter. You must attempt to obtain the results of supplemental testing required under §610.40(e) of this chapter prior to notifying a donor of the deferral. If notification occurs prior to receipt of such results, you must also notify a deferred donor of the results of the supplemental testing. You must notify a donor as described in paragraph (b) of this section.
- (b) Content of notification. You must provide the following information to a donor deferred or determined not to be suitable as a donor as described in paragraph (a) of this section:
- (1) That the donor is deferred or determined not to be suitable for donation and the reason for that decision;

- (2) Where appropriate, the types of donation of blood or blood components that the donor should not donate in the future:
- (3) Where applicable, the results of tests for evidence of infection due to communicable disease agent(s) that were a basis for deferral under §610.41 of this chapter, including results of supplemental (i.e., additional, more specific) tests as required in §610.40(e) of this chapter; and,
- (4) Where appropriate, information concerning medical followup and counseling.
- (c) Time period for notification. You must make reasonable attempts to notify the donor within 8 weeks after determining that the donor is deferred or determined not to be suitable for donation as described in paragraph (a) of this section. You must document that you have successfully notified the donor or when you are unsuccessful that you have made reasonable attempts to notify the donor.
- (d) Autologous donors. (1) You also must provide the following information to the referring physician of an autologous donor who is deferred based on the results of tests for evidence of infection with a communicable disease agent(s) as described in paragraph (a) of this section:
- (i) Information that the autologous donor is deferred based on the results of tests for evidence of infection due to communicable disease agent(s), as required under §610.41 of this chapter, and the reason for that decision;
- (ii) Where appropriate, the types of donation of blood or blood components that the autologous donor should not donate in the future; and
- (iii) The results of tests for evidence of infection due to communicable disease agent(s), that were a basis for deferral under §610.41 of this chapter, including results of supplemental (i.e., additional, more specific) tests as required in §610.40(e) of this chapter.
- (2) You must make reasonable attempts to notify the autologous donor's referring physician within 8 weeks after determining that the autologous donor is deferred as described in paragraph (a) of this section.

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You must document that you have successfully notified the autologous donor's referring physician or when you are unsuccessful that you have made reasonable attempts to notify the physician.

PART 640—ADDITIONAL STAND-ARDS FOR HUMAN BLOOD AND BLOOD PRODUCTS

Subpart A—Whole Blood

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640.3	Suitability of donor.
640.4	Collection of the blood.
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Subpart B—Red Blood Cells

640.10	Red Blood Cells.
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640.12	Suitability of donor.
640.13	Collection of the blood.
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Subpart C—Platelets

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Subpart D—Plasma

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subpair E [Reserved]

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640.51	Suitability of donors.
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Subpart G—Source Plasma

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640.60 Source Plasma.
640.61 Informed consent.
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640.62	Medical supervision.	
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640.63 Suitability of donor.

640.64 Collection of blood for Source Plasma.

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640.65 Plasmapheresis.

640.66 Immunization of donors.

640.67 Laboratory tests.

640.68 Processing.

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640.70 Labeling.

640.71 Manufacturing responsibility.

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640.73 Reporting of fatal donor reactions.

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Subpart H—Albumin (Human)

640.80 Albumin (Human).

640.81 Processing.

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Subpart J—Immune Globulin (Human)

640.100 Immune Globulin (Human).

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640.102 Manufacture of Immune Globulin (Human).

640.103 The final product.

640.104 Potency.

Subpart K [Reserved]

Subpart L—Alternative Procedures

640.120 Alternative procedures.

AUTHORITY: 21 U.S.C. 321, 351, 352, 353, 355, 360, 371; 42 U.S.C. 216, 262, 263, 263a, 264.

SOURCE: 38 FR 32089, Nov. 20, 1973, unless otherwise noted.

CROSS REFERENCES: For U.S. Customs Service regulations relating to viruses, serums, and toxins, see 19 CFR 12.21–12.23. For U.S. Postal Service regulations relating to the admissibility to the United States mails see parts 124 and 125 of the Domestic Mail Manual, that is incorporated by reference in 39 CFR part 111.

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